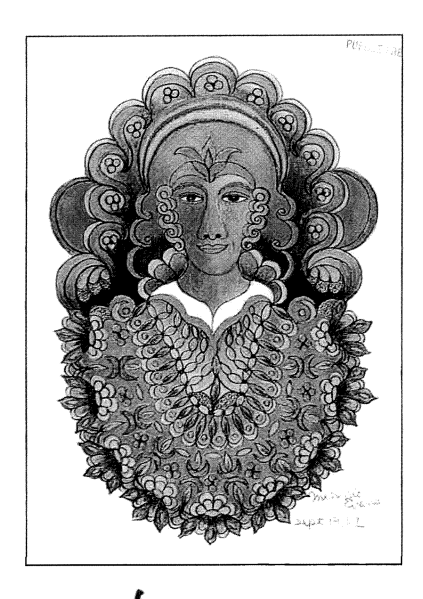
# EXHIBIT A

November 13, 2002





\*\*\*\*\*\*\*CR LOT 0003A\*\*C-006 0001 STAMFORD CT 06901-3431 986 051 Madhaldallamillallalalallaidllaadlallall 628

The most serious risk associated with opioids, including OxyContin<sup>s</sup>, is respiratory depression. Common opioid side effects are constipation, nausea, sedation, dizziness, vomiting, pruritus, headache, dry mouth, sweating and weakness.

OxyContin<sup>®</sup> is contraindicated in patients with known hypersensitivity to oxycodone, or in any situation where opioids are contraindicated.

Please see Contraindications section in package insert.

Purdue is firmly committed to maintaining the highest standards of marketing practices in the industry while continuing to advance the proper treatment of pain in America. If Purdue's marketing and sales practices fail to meet this standard, we urge you to contact us at 1-888-690-9211.



(OXYCODONE HCI CONTROLLED-RELEASE) TABLETS

Please read brief summary of prescribing information including boxed warning on reverse side.

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A7087-F5

FUR-40009278



\*80 mg and 160 mg for use in opioid-tolerant patients only

OxyContin is an opioid agonist and a Schedule II controlled substance with an abuse liability similar to morphine.

Oxycodone can be abused in a manner similar to other opioid agonists, legal or illicit. This should be considered when prescribing or dispensing OxyContin in situations where the physician or pharmacist is concerned about an increased risk of misuse, abuse, or diversion.

OxyContin Tablets are a controlled-release oral formulation of oxycodone hydrochlo-ride indicated for the management of moderate to severe pain when a continuous around-the-clock analgesic is needed for an extended period of time.

### OxyContin Tablets are NOT intended for use as a prn analoesic.

OxyContin 80 mg and 160 mg Tablets ARE FOR USE IN OPIOID-TOLERANT PATIENTS ONLY. These tablet strengths may cause fatal respiratory depression when administered to patients not previously exposed to opioids.

OXYCONIIN TABLETS ARE TO BE SWALLOWED WHOLE AND ARE NOT TO BE BRO-Ken, Chewed, or Crushed. Taking Broken, Chewed, or Crushed Oxyconiin Tablets Leads to rapid release and absorption of a potentially fatal Dose of Oxyco

OxyContin\* (oxycodone hydrochloride controlled-release) Tablets are an opioid analgesic supplied in 10 mg, 20 mg, 40 mg, 80 mg, and 160 mg tablet strengths for oral administration.

### INDICATIONS AND USAGE

ish<sup>®</sup> Tablets are a controlled-release oral formulation of oxycodone hydrochloride indicated for the mant of moderate to severe pain when a continuous, around-the-clock analgestic is needed for an d period of time.

### OxyContin is NOT intended for use as a orn analoesic.

Conjugation in Not institute on use see pin relegacy.

Physicians should individualize treatment in every case, initiating therapy at the appropriate point along a progression from non-opioid analysiscs, such as non-steroidal ant-inflammatory drops and acation, the microphen to policia in palan of plan innangement such as outlined by the World Health Chipart Saloti, the Agency for Healthcare Research and Quality (formerly known as the Agency for Health Case Policy and Research), the Tederation of State Medical Boards Model Cuidelines, or the American Palin Society.

research), the Federation of State Medical Boards Model Cidelines, or the American Pain Society.

Op/Confin Is not indicated for pain in the immediate postoperative period (the first 12-24 hours following surgery), or if the pain is mild, or not expected to persist for an extended persiod of time. Op/Confin Is only indicated for postoperative use if the patient is afready receiving the drug prior to surgery or if the post-persitive pain is expected to he most certain to severe and presist for an extended persiod of time. Purplications should individualize treatment, moving from parenteral to oral analysis as appropriate. (See American Para Society quidelines.)

OxCordin' is contraindicated in patients with known hypersensitivity to oxycodone, or in any situation where opolicis are contraindicated. This includes patients with significant respiratory depression (in unmonitored settings or the absence of resuscitative equipment), and petiants with acute or severe bronchial safety in the operation of t

### WARNINGS

WAHNINIS

OVYCONTIN TABLETS ARE TO BE SWALLOWED WHOLE, AND ARE NOT TO BE BROKEN, CHEWED OR GUISHED. TAKING BROKEN, CHEWED OR GUISHED. TAKING BROKEN, CHEWED OR GUISHED DAYCONTIN TABLETS LEADS TO RAPID RELEASE AND ASSORPTION OF A POTENTIALLY FATAL DOSE OF OXYCODONE.

OxyConin 80 mg and 160 mg Tablets ARE FOR USE IN OPTIOL-TOLERANT PATIENTS ONLY These tablet strengths may cause tatal respiratory depression when administered to patients not previously exposed to optoids.

exposed to opioids.

OxyContin 80 mg and 160 mg Tablels are for use only in opioid-tolerant patients requiring daily oxycodone squivalent dosages of 160 mg or more for the 80 mg tablet and 220 mg or more for the 180 mg tablet. Care should be taken in the prescribing of these tablet strengths. Patients should be instructed against use by individuals other than the patient for whom It was prescribed, as such inappropriate use may have severe medical consequences, including death.

Misuse, Abuse and Diversion of Opioids

Oxycodone is an opicid agonist of the morphine-type. Such drugs are sought by drug abusers and peo-ple with addiction disorders and are subject to criminal diversion.

pie with addiction osorders and are subject to criminal diversion.

Opvocatione can be abused in a manner similar to other opidia gonists, legal or illicit. This should be considered when prescribing or dispensing OxyCordin in situations where the physician or pharmacist is concerned about an increased risk of milisace, abuse, or diversion, snorting, or injecting the dissovery product. These practices will result in the uncontrolled delivery of the opidical and poss a significant risk to the abuser that could result in oversion death (see WARNINGS and DRUG ABUSE AND ADDICTION).

the abuser that could result in overgoes and oeath (see WARHINIAS and DHOL ABUSE AND ADULE TIMP).

Concerns about abuse, addiction, and diversion should not prevent the proper management of pain. The
development of addiction to opioid analgesics in properly managed patients with pain has been reported
to be rare. However, data are not available to establish the true incidence of addiction in chronic pain patients. Healthcare professionals should contact their State Professional Licensing Board, or State Controlled Substances Authority for information on how to prevent and detect abuse or diversion of this product.

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# DRUG ABUSE AND ADDICTION

OxyContin' is a mu-agonist opinid with an abuse liability similar to morphine and is a Schedule II con-troiled substance. Oxycodone, like morphine and other opioids used in analgesia, can be abused and is subject to criminal diversion.

Drug addiction is characterized by compulsive use, use for non-medical purposes, and continued use despite harm or risk of harm. Drug addiction is a treatable disease, utilizing a mutit-disciplinary approach, but relapse

Drug-seeking' behavior is very common in addicts and drug abusers. Drug-seeking tactics include emergency calls or visits are the end of office hours, refusal to undergo appropriate camilation, test-ing or reteral, regarded loss of prescriptions, tampering with prescriptions and refuciacies to provide prior medical records or contact information for other realing physicians). 'Doctor shopping to obtain addi-tional prescriptions is common among drug abusers and people suffering from unfreated addictions.

bonal prescriptions is common among drug abusers and people suffering from uniteral ablocition. Abuse and addiction are spanare and siliciant from physical dependence and otherance. Physicians should be aware that addiction may not be accompanied by concurrent tolerance and symptoms of physical dependence in all addicts. In addiction, allow of points can cover in the absence of three addiction and scharacterized by misuse for non-medical purposes, other in combination with other psychoarbine substances. Oncycomin', like other opticis. Is as been othered for non-medical sus. Careful foront Aespiring of prescriping information, including quantity, frequency, and renewal requests is strongly advised.

Proper assessment of the patient, proper prescribing practices, periodic re-evaluation of therapy, and proper dis-pensing and storage are appropriate measures that help to limit abuse of opioid drugs.

persorg and sorage are appropriate measures that free to initial cause of option drugs. On/Contin consists of a dual-polymer matrix, intended for or all use only. Abuse of the crushed tablet posse a hazard of overdose and death. This risk is increased with concurrent abuse of alcohol and other substances. With parenteral abuse, the hablet explicitle, especially late, can be expected for all in local lissue necrosis, infection, pulmonary granulomas, and increased risk of endocarditis and valvular heart injury. Perenteral drug abuse is commonly associated with transmission of infectious diseases such as hepatitis and HIV.

# Respiratory Depression

Respiratory depression is the chief hazard from oxycodone, the active ingredient in OxyContin\*, as with all opi-oid agonists. Respiratory depression is a particular problem in elderly or debilitated patients, usually following large initial coses in non-tolerant patients, or when opoids are given in conjunction with other agents that depress

respirators. Devycobne should be used with externe caution in patients with significant chronic obstructive pulmonary disease or or pulmonary, and in patients having a substantially decreased respiratory reserve, hypoxia, hypercaria, or pre-existing respiratory depression, it usual patients, were usual herazellut dosses of oxylogical patients, and produce the properties of special highest the description of the patients of the

The respiratory depressant effects of opioids include carbon dioxide retention and secondary elevation of ceebiospinal fluid pressure, and may be markedly exaggerated in the presence of head injury, intract-nal lesions, or divide sources of pre-eleviting increased intractantal pressure. Opcoding produces effects on pupilary response and consciousness which may obscure neurologic signs of further increases in intracra-nal pressure in patients with head injury.

OxyContin may cause severe hypotension. There is an added risk to individuals whose ability to maintain blood

pressure has been compromised by a degleted bod volume, or after countert administration will as phenothazens or other agents which compromise vesorion force. Oxycodore may produce or hos sion in ambitatory patients. Oxycodore, like all opioid analysists of the morphine-type, should be awdit caution to peters in circulatory shock, since vascollation produced by the drug may further data output and blood pressure.

PREDERITIONS

### PRECAUTIONS

### General

Opioid analgesics have a narrow therapeutic index in certain patient populations, especially when com-bined with CNS depressant drugs, and should be reserved for cases where the benefits of opioid analge-sia outweigh the known risks of respiratory depression, aftered mental state, and postural hypotension.

Use of OxCortin 1s associated with increased potential risks and should be used only with caution in the tollowing conditions acute alcoholism, adenocortical insufficiency (e.g., Addison's disease). CNS depres-sion or come, definity memers, debilitated playerts, kpylosocious associated with respiratory depression, impacted an or hypothyroidism; prostatic hyperropty or urethral shirbure; severe impairment of hepatic, pul-monary or real function; and tode; psychosis.

initially of initial miniouri, and tool psychosis. The administration of oxycodone may obscure the diagnosis or clinical course in patients with acute abdom inal conditions. Dxycodone may aggravate convulsions in patients with convulsive disorders, and all op-olds may induce or aggravate seizures in some clinical settings.

# Interactions with other CNS Depressants

Day Contan should be used with custom and started in a reduced dosage ( $^{1}/_{2}$  to  $^{1}/_{2}$  of the usual dosage) in patients who are concurrently receiving other central nervous system depressaris motioning sectaines or hymorics, general ansethetics, phenoinizanies, other tranquizers, and alcohol. Interactive effects resulting an respiratory depression, hypotension, profound sedation, or coma may result if these drugs are taken in combination with the usual doses of OxyCortin.

taxen in commonation with usual obsess or usycoreum. Interactions with wixed Aponist'Antangonist Opioid Analgesics Aponds'artagonist analgesics (i.e., pertazonien, enabuphine, and butorphanol) should be administered with action to a patient with on has received or is receiving a course of therapy with a pure opioid agonist anal-gesic such as oxycodone. In this shaution, mode agonisty'artagonist analgesics may reduce the analgesic effect of oxycodone and/or may precipitate withdrawal symptoms in these patients.

# Ambulatory Surgery and Postoperative Use

Amoustory Surgery and Postoperative Use
OxyContin is not indicated for pre-emptive analgesia (administration pre-operatively for the management
opsosperative pain).
OxyContin is not indicated for pain in the immediate postoperative period (the first 12 to 24 hours following surgery) for patients not previously taking the drug, because its safety in this setting has not
been established.

OxyContin is not indicated for pain in the postoperative period if the pain is mild or not expected to per-sist for an extended period of time.

NAME TO AN EXEMBED PRIOR OF MINE.

Opp.Contin's los only indicated for postoperative use if the patient is already receiving the drug prior to surgery or if the postoperative pain is expected to be moderate to severe and persist for an extended period of time. Physicians should individualize treatment, moving from parenteral to oral analgesics as appropriate (See American Pain Society guidelines).

Patients who are already receiving OwComin Tablets as part of ongoing analgesic thraziny may be safely continued on the drug if appropriate disage adjustments are made considering the procedure, other drugs given and the importance of the processing three processing three processing of the procedure, other drugs given and the importance of the processing three processing three

DOSAGE AND ADMINISTRATION).

One Confined and their morphine-like opioids have been shown to decrease bowel motility, lieus is a common postoperative complication, especially after intra-abdominal surgery with opioid analgesia. Caution should be taken to motific for decreased bowel motility in postoperative patients receiving opioids. Standard supportive therapy should be implemented. Use in Pancreatic/Biliary Tract Disease

Oxycodone may cause spasm of the sphincter of Oddi and should be used with caution in patients with billiary tract disease, including acute pancreatitis. Opioids like oxycodone may cause increases in the serum amylase level.

Tolerance is the need for increasing doses of opioids to maintain a defined effect such as analyses (in the absence of desase progression or other external factors). Physical dependence is manifested by withdrawal symptoms after abrupt discornization of a drug or upon administration of an antagonist. Physical dependence and to-erance are not unusual during chronic opioid therapy.

the opioid abstinence or withdrawal syndrome is characterized by some or all of the following: restless-ness, lacrimation, rhinorrhea, yawning, perspiration, chills, myalqia, and mydriasis. Other symptoms also may develop, including: intability, aroxiety, backache, joint pain, weakness, abdominal cramps, insomnia, nausea, ancrexia, vomitimg, diarrinea, or increased blood pressure, respiratory rate, or heart rate.

# In general, opioids should not be abruptly discontinued.

### Information for Patients/Caregivers

information by Taulery and great of the follow-to clinically advisable, patients receiving OxyContin Tablets or their caregivers should be given the follow-ing information by the physician, nurse, pharmacist, or caregiver:

1. Patients should be aware that OxyContin Tablets contain oxycodone, which is a morphine-like substance.

- Patients should be advised that DoyContin Tablets were designed to work properly only if swallowed whole. OxyContin Tablets will release all their contents at once if broken, chewed, or crushed, resulting in a risk of fatal overdose.
- Patients should be advised to report episodes of breakthrough pain and adverse experiences occurring during therapy, individualization of dosage is essential to make optimal use of this medication.
   Patients should be advised not to adjust the dose of OxyContin' without consulting the prescribing pro-

- Tessions.

  Features should be advised that Op/Contin may impair mental and/or physical ability required for the per-formance of potentially hazardous tasks (e.g., driving, operating hasy machinery).

  Patients should not combine Op/Contin with alcohol or other central nervous system depressants (sleep acids, barquitagers) goods by the uniform of death continuing physician, because temperous additive effects

  Woman of childrening potential with o become, or are planning to become, preparat should be advised to council the driving physician preparation.
- Patients should be advised that OxyContin is a potential drug of abuse. They should protect it from theft and it should never be given to anyone other than the individual for whom it was prescribed.
- Patients should be advised that they may pass empty matrix 'ghosts' (tablets) via colostomy or in the stool, and that this is of no concern since the active medication has already been absorbed.
- stoot, and that this is of no concern since the active medication has already been absorbed.

  10. Patients should be advised that if they have been receiving treatment with Opcontrin former than a few weeks and cessation of therapy is indicated, it may be appropriate to taper the Opcontrin dose, rather than aburphy discontinual, due to the risk of precipitating withdrawed symptoms. Their physician, can provide a dose scriedule to accomplish a gradual discontinuation of the medication.

  11. Patients should be instructed to keep DoyCorthin in a secure place out of the reach of children. When OyCorthin is no longer receded, the unusued tables should be destroyed by flushing down the toilet.

  12. The received in the control of the stood of the control of the contro

OxyCorthis is an option with no approved use in the management of addictive disorders. Its proper usage in individuals with drug or alcohol dependence, either active or in remission, is for the management of pain requiring opioid analgesia.

# **Drug-Drug Interactions**

uring-uring interactions. Opioid analysiss, including (bxyContin\*, may enhance the neuromuscular blocking action of skeletal mus-cle relearants and produce an increased degree of respiratory depression. Opyoconde is Berabbicated in part to opyomophone via cycloxorane P450 206. While this pathway may be blocked by a variety of drugs (e.g., certain cardiovascular drugs including amiodarone and quilidine as well as polycyclic artidepressants), such bloosade has not yet been shown to be of linical significance with this agent. Clinicians should be aware of this possible interaction, however.

# Use with CNS Depressants

Use with CNB Depressants 
Opcontrol. Nee all opioid analysiss, should be started at 1/2 to 1/2 of the usual dosage in patients who are concurrently receiving other central nervous system depressants including setatives or hyporotics, pere all aresthetics, phenothazines, centrally acting anti-emetics, tranquillors, and actional because respiratory depression, hypotension, and profound setation or coma may result. No specific interaction between ocyclobre and monorarine coldates inhibitors has been observed, but caution in the use of any opioid in patients taking his class of drugs is appropriate.

# Carcinogenesis, Mutagenesis, Impairment of Fertility

# Studies of oxycodone to evaluate its carcinogenic potential have not been conducted

sources or usystemson to excusive its carcinogenic potential have not been conducted. Opporation was not mulagenic in the following assays: Ames Satmonetia and E coil test with and without metabolic activation at disess of up to 5000  $\mu$ g, chromosomal aberration test in human hymbrough in the absence of metabolic activation at disess of up to 1500  $\mu$ g/m and with anxietation 48 hours after exposure at doses of up to 1500  $\mu$ g/m., and in the in two bone merrow micronucleus test in mixed state exposure at doses of up to 1500  $\mu$ g/m., and in the in two bone merrow micronucleus test in mixed state exposure at doses of up to 1500  $\mu$ g/m., and in the in who bone merrow micronucleus test in mixed assay in the presence of metabolic activation in the human chromosomal aberation test (at greator than equal to 1250  $\mu$ g/m). It 27 but not real 8 hours of exposure and in the mouse hymphoma assay at doses of 50  $\mu$ g/mL or greater with metabolic activation and at 400  $\mu$ g/mL or greater without metabolic activation.

Paradogenic Effects — Category B: Reproduction studies have been performed in rats and rabbits by oral administration at classes up to 6 mpkg and 125 mpkg, respectively. These does are 3 and 46 times a human date of 160 mpkg, based on mpkg bases. The results during the results of the result

CopyContin' is not recommended for use in women during and immediately prior to labor and delivery because or prior to produce any cause respiratory depression in the newborn. Neonates whose mothers have been taking oxycodone chronically may exhibit respiratory depression and/or withdrawal symptoms, either at birth models in the amount of the production of the productio

normal mouters Low concentrations of oxycodone have been detected in breast milk. Withdrawal symptoms can occur in breast-leading infants when maternal administration of an opioid analgesic is stopped. Ordinarily, nursing should not be undertaken while a patient is receiving OxyContin because of the possibility of setation and/or respiratory degression in the infant.

Geratario Use
In controlled pharmacokinetic studies in elderly subjects (greater than 65 years) the clearance of oxyodome appeared to be slightly reduced. Compand to young adults, the plasma concentrations of oxyodome experance of the slightly reduced. Compand to young adults, the plasma concentrations of oxyodome version creased appromisedly 155. Of the trail number of subjects (445) in cincilar studies of Oxyodomit. 146 (33.3%) were age 65 and older (including those age 75 and older) while 40 (9.0%), were age 75 and older of cincilar studies of the appropriate inlaints on the train of cost trailard, no unitivosar or unappeaded side effects were seen in the elderly patients who received Oxyodomin. Thus, the usual dosea and dosing printer size are propried for three patients. As with all politics, the sturing dose stood be reduced to 1/2 to 1/2 of the usual dosage in debitated, non-tolerant patients. Respiratory depression is the chief hazard in elderly or debitated patients, usually following large initial doses in on-tolerant patients, or when opioids are given in conjunction with other agents that depress respiration.

### Laboratory Monitoring

Due to the those drange of plasma concentrations seen in clinical populations, the varying degrees of pain and the development of tolerance, plasma oxycodone measurements are usually not helpful in clinical man agement. Plasma concentrations of the active drug substance may be of value in selected, unusual or com

### Hepatic Impairment

study of OxyContin in patients with hepatic impairment indicates greater plasma concentrations than those ifth normal function. The initiation of therapy at  $^{1}/_{3}$  to  $^{1}/_{2}$  the usual doses and careful dose titration is

### Renal (mpairment

In patients with enail impairment, as evidenced by decreased creatinine clearance (<60 mL/min), the con-centrations of oxycodorie in the plasma are approximately 50% higher than in subjects with normal renal function. Dose initiation should follow a conservative approach. Desegres should be adjusted according to the clinical standard. **Gender Differences** 

In pharmacoloinetic studies, oploid-naive females demonstrate up to 25% higher average plasma con-centrations and greater frequency of hybrid oploid adverse event blam males, even after adjustment for body weight. The clinical relevance of adfreence of this magnitude is low for a drug intended for ic usage at individualized dosages, and there was no male/female difference detected for efficacy or adverse

### ADVERSE REACTIONS

The safety of Do-Contin' was evaluated in double-blind clinical trials involving 713 patients with moderate to severe pain of various etiologies. In open-label studies of cancer pain, 187 patients received OxyContin in total child yoses ranging from 20 mg to 640 mg per day. The average total daily dose was approximately 105 mg per day.

oxyconia in rusus tusing obests staging into 1, mig to down tip, excitate the transpire of initial use are those observed with other point an algorithms and several reactions which may be associated with OxyContin Tablet the transpire in clinical use are those observed with other quoted analysistic, including respiratory depression, aporta, respiratory arrest, and (to an even issuer degree) circulations depression, hypotension, or shock (see OVERIOSAGE). The non-serious eventures exercises on existence of the transpire of the OxyContin are hypical oxidiol side. The non-serious exercise continued to the continued of the continued and some degree of the continued on the continued of the continued and some degree of the continued on the continued of the continued of the continued of the continued and some degree of the continued on the continued of the continued o

apy is continuous and some organs on what immediate-release oxycodone and placebo revealed a similar adverse event profile between OxyCortin with immediate-release oxycodone. The most common adverse events (>5%) reported by patients at least once during therapy were:

	OxyContin (n=227) (%)	Immediate- Release (n=225) (%)	Placebo (n=45) (%)	
Constipation	(23)	(26)	(7)	
Nausea	(23)	(27)	(11	
Somnolence	(23)	(24)	(4)	
Dizziness	(13)	(16)	. (9)	
Pruritus	(13)	(12)	(2)	
Vomiting	(12)	(14)	(7)	
Headache	(7)	. (8)	(7)	
Dry Mouth	(6)	. (7)	(2)	
Asthenia	(6)	(7)		
Sweating	(5)	(6)	(2)	

The following adverse experiences were reported in DayContin\*-treated patients with an incidence between 1% and 5%, in descending order of frequency they were ancreak, nervousness, insomnia, tever confusion, diorrhea, abdominal pain, dyspepsia, rash, anxiety, euplinoi, dyspera, postural hypotension, chills, twitching, gastitis, abnormal dreams, thought abnormalities, and hiccups.

The following adverse reactions occurred in less than 1% of patients involved in clinical trials or were reported in postmarketing experience.

General: accidental injury, chest pain, facial edema, malaise, neck pain, pain

Cardiovascular: migraine, syncope, vasodilation, ST depression Digestive: dysphagia, eructation, flatulence, gastrointestinal disorder, increased appetite, nausea and vomiting, stomatitis, lieus

vornting, stomatitis, letus:
Hemic and Lymphatic imprisadenopalhy
Metabolic and Nutritionat delitydration, edema, hyponatremia, peripheral edema, syndrome of inappropridia antiduretic hormone secretion, thirst
Nervous: ahnormal gait, agitation, ammesia, depersonalization, depression, emotional lability, hallucination, hyperinesia, hypesthesia, hypotonia, malaise, parestinesia, setzieres, seech disorder, stupor, firmitus, hemor, verigo, withdrawal syndrome with or without sectures

Respiratory: cough increased, pharyngitis, voice afteration

Special Senses: abnormal vision, taste perversion norrhea, decreased libido, dysuria, hematuria, impotence, polyuria, urinary retention, uri-

Acute overdosage with oxycodone can be manifested by respiratory depression, somnolence progress-ing to stupor or coma, skeletal muscle flaccidity, cold and clammy skin, constricted pupils, bradycardia, hypotension, and death.

represension, and usual.

Deaths due to overdose have been reported with abuse and misuse of OxyContin\*, by ingesting, inhaling or injecting the crushed tablets. Review of case reports has indicated that the risk of fatal overdose is fur ther increased when OxyContin is abused concurrently with alcohol or other CNS depressants, including

ourer upunus.
In the teatment of oxycodone overdosage, primary attention should be given to the re-establishment of a patent airway and institution of assisted or controlled ventilation. Supportive measures (including oxyen and vasopressors) should be employed in the management of circulatory slock and parimonary sterma accompanying overdose as indicated. Cardac arrest or arrhythmias may require cardiac massage or delib-

rillation. The pure opioid antagonists such as nationone or nalmetene are specific antidotes against respiratory depres-sion from opioid overdose. Opioid antagonists should not be administered in the absence of finicials tra-niciant respiratory or circulatory depression secondary to polycordone overdose, in patients who are physically dependent on any opioid agrinatin reliability. Opioid finitial respiratory are proprietable and aprincial reliability of complete reversal of opioid effects many perceptable a more on pressule depresentation and the proprietation of the

# naging Expected Opioid Adverse Experiences

managing expected upons waverse experiences
Most patients receiving opidist, sepsicially those who are opioid-naive, will experience side effects. Frequently the side effects from OnyCordin are transient, but may require evaluation and management. Advancements such as consciptional should be articipated and treation agressively and prophylactically with a standard and treation and treating and the standard and treating and the standard and treating and treating and the standard and the standard

Other opioid-related side effects such as sedation and nausea are usually self-limited and often do not per-sist beyond the first lev days. If nausea persists and is unacceptable to the patient, treatment with aritemet-ics or other modalities may relieve these symptoms and should be considered. Patients receiving looyConfrir may asks a mintact matric (1905) if the stood or via colostomy. These ghosts contain little or no residual polycodone and are of no clinical consequence.

NAPLET AND HAMULING

OyContin Tablets are solid dosage forms that contain coycodone which is a controlled substance. Like morphine, oycodone is controlled under Schedule II of the Controlled Substances Act.

OxContin has been targeted for theft and diversion by criminals. Healthcare professionals should contact their site Professional Libersing Board or Sales Controlled Substances Authority for information on how to prevent and detect abuse or diversion of this product.

Single 2015 CTPS experience partities the hashes 15°-2010\* (50-28-5).

Store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F).

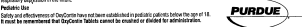
Healthcare professionals can telephone Purdue Pharma's Medical Services Department (1-888-726-7535) for information on this product.

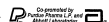
# DEA Order Form Required.

e Pharma L.P., Stamford, CT 06901-3431

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U.S. Patent Numbers 4,861,598; 4,970,075; 5,266,331; 5,508,042; 5,549,912; and 5,656,295





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